

Global metagenomic survey reveals taxonomic "blind spots" in geothermal springs

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Abstract

Analysis of the increasing wealth of metagenomic data collected from diverse environments can lead to the discovery of novel branches on the tree of life. Here we analyse 5.2 Tb of metagenomic data collected globally to discover a novel bacterial phylum ('Candidatus Kryptonia') found exclusively in high-temperature pH-neutral geothermal springs. This lineage had remained hidden as a taxonomic "blind spot" due to mismatches in the primers commonly used for ribosomal gene surveys. Genome reconstruction from metagenomic data combined with single-cell genomics results in several high-quality genomes representing four genera from the new phylum. Metabolic reconstruction indicates a heterotrophic lifestyle with conspicuous nutritional deficiencies, suggesting the need for metabolic complementarity with other microbes. Co-occurrence patterns identifies a number of putative partners, including an uncultured Armatimonadetes lineage. The discovery of Kryptonia within previously studied geothermal springs underscores the importance of globally sampled metagenomic data in detection of microbial novelty, and highlights the extraordinary diversity of microbial life still awaiting discovery.

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Molecular environmental surveys have provided a sizeable snapshot of microbial phylogenetic diversity. Sequencing of small subunit ribosomal RNA (SSU rRNA) genes directly from the environment has expanded the known microbial tree of life from Woese's original twelve phyla to more than 70 bacterial phyla^{1,2}. Advances in cultivation-independent methods for examining uncultured microbes, including single-cell genomics and deep sequencing of environmental samples, have begun yielding complete or near-complete genomes from many novel lineages³⁻¹⁰. These approaches have already led to the recovery of genomic information from a wealth of candidate lineages (phylogenetic lineages for which a cultured representative is not available), notably the Lokiarchaeota¹¹, Pacearchaeota and Woesearchaeota¹⁰, and members of the Candidate Phyla Radiation³. These lineages, previously recognized only through SSU rRNA data and residing in poorly sampled habitats, are providing a more complete topology of the tree of life.

More recently, it has been suggested that a wealth of novel bacterial and archaeal clades exist that are systematically under-represented (the 'rare biosphere') or missed altogether in classical surveys, leaving significant taxonomic "blind spots" Compared to many of the proposed candidate phyla for which SSU rRNA gene information exists, these taxonomic "blind spots" are uncharted lineages with potentially important ecological and evolutionary implications. Further, these lineages may be highly abundant and hold important metabolic or functional roles within the community, yet have been overlooked thus far in ecological surveys. Metagenome sequencing is uniquely suited for uncovering taxonomic "blind spots" since it does not suffer from

biases introduced during PCR amplification, and has limitations only with insufficient resolution of minor populations within a community. However, an exploration of the complete compendium of available metagenomic sequences for the presence of taxonomic "blind spots" has yet to be performed¹³. Here, we report the results of large-scale mining of metagenomic data and single cell genomics, which led to the discovery of a new bacterial phylum in geographically distinct geothermal springs.

Results

Identification of a novel bacterial candidate phylum

To cast a global net for the discovery of novel microbial lineages in the absence of biases introduced via PCR amplicon-based surveys, we collected long assembled contigs (≥ 100 kbp) from a comprehensive collection of 4,290 metagenomic datasets available through the Integrated Microbial Genomes with Microbiome Samples (IMG/M), a database containing a total of more than 5 Tb of sequence data¹⁴. From these data, 31,955 assembled contigs were identified and 744 contigs were further selected that contained SSU rRNA gene fragments greater than 100 bp (Fig. 1A). The SSU rRNA gene sequences were then aligned and phylogenetically placed on a reference tree consisting of high-quality SSU rRNA sequences from Bacteria and Archaea¹5,¹6.

Exploration of the constructed SSU rRNA tree for novel phylogenetic branches led to the identification of a distinct lineage consisting of a full-length SSU rRNA sequence. A subsequent search against all assembled metagenomic data identified three additional full-length SSU rRNA sequences. The four SSU rRNA gene sequences were from four geographically distant, high-temperature, pH-neutral, geothermal springs in North

America and Asia (Fig. 1). These sequences shared an average 97.4% identity (± 1.97% s.d.), and showed a maximum identity of only 83% to SSU rRNA genes (such as the one in GenBank ID: AP011715) in NCBI's Non-Redundant (NR) database. In line with the notion of taxonomic "blind spots" 12, a comparison of 'universal' SSU rRNA primer sets typically used for full-length and hypervariable region amplification with the four novel sequences indicated numerous mismatches, explaining why members of this lineage likely eluded detection in previous microbial diversity surveys (Supplementary Fig. 1; Supplementary Table 1).

Phylogenetic analysis of the four SSU rRNA genes placed the newly discovered lineage into a monophyletic branch within the *Fibrobacteres-Chlorobi-Bacteroidetes* (FCB) superphylum^{9,17} (Supplementary Fig. 2). Based on suggested thresholds for SSU rRNA sequence identity to distinguish new phyla^{2,18}, we propose that this lineage represents a new bacterial candidate phylum (Supplementary Table 2).

Comparative genomics and cell morphology of novel FCB lineage

Reassembly of the metagenomic data combined with tetranucleotide-based binning methods using the initial contigs containing the SSU rRNA genes yielded near-complete recovery of four distinct genomes, each from one of the four spring samples (Supplementary Fig. 3; Supplementary Table 3). Phylogenetic analysis of conserved marker genes supported its placement as a sister phylum to the *Ignavibacteria* with 100% bootstrap support (Fig. 2A; Supplementary Fig. 4). Three of the genomes reconstructed from metagenomes (GFMs) from Dewar Creek Spring, Canada¹⁹, Great Boiling Spring, Nevada^{20,21}, and Gongxiaoshe pool, Yunnan Province, China²² had an

average 95.8% estimated coverage, while the genome from Jinze pool, Yunnan Province, China²² had a lower estimated coverage of 68% (Supplementary Table 4). The high genomic sequence coverage across the four metagenomes (average 31.2x coverage; Supplementary Table 3) suggested that this novel lineage might exist at sufficient cell abundance to be captured by single cell technology. We therefore employed high-throughput single-cell isolation, whole-genome amplification and SSU rRNA screening of single amplified genomes (SAGs) in search for the novel lineage (Fig. 1). We successfully recovered a total of 18 SAGs from three of the four samples, corresponding to the novel phylum-level clade with an estimated average genome completeness of 67.2% (± 20.1 s.d.) (Supplementary Table 3). We designate this new candidate phylum 'Candidatus Kryptonia,' from the Greek word 'krupton' meaning hidden or secret since it has hitherto eluded detection due to SSU rRNA primer biases (Supplementary Table 4).

The average nucleotide identity (ANI) based metric, Microbial Species Identifier (MiSI), was used to compare the four 'Ca. Kryptonia' genomes reconstructed from metagenomes (GFMs) and the 18 SAGs²³. This analysis revealed that almost all of the genotypes extracted from the same sample belonged to a single species (Supplementary Table 5). For example, the GFM reconstructed from Dewar Creek ('Ca. Kryptonium thompsoni' JGI-4) and the thirteen SAGs ('Ca. Kryptonium thompsoni' JGI-5 – JGI-17) collected from the same site shared an ANI of 99.67% (± 0.15 s.d.) and represent a single coherent species²³. A single exception to the above observations was the recovery of a divergent 'Ca. Kryptonia' SAG ('Ca. Chrysopegis kryptomonas' JGI-23) from the Jinze pool, Yunnan Province, China representing a population distinct from

the other two SAGs recovered from this site ('Ca. Kryptobacter tengchongensis' JGI-24 and JGI-25) (Supplementary Table 5). Across the four geothermal springs, the GFMs and SAGs collectively share average ANIs of only 78.86% (± 1.42 s.d.), suggesting that they represent different genera of 'Ca. Kryptonia'. Further support for genus-level designations is evident from nuanced functional and metabolic differences across the genomes, as described below.

In addition to recovering single cells of *Ca.* Kryptonia for genome amplification, we designed a SSU rRNA-targeted fluorescence *in situ* hybridization (FISH) probe to visualize cell morphology (Fig. 2B). The targeted *Ca.* Kryptonia cells appeared filamentous, and exhibited morphological heterogeneity ranging from short to elongated filaments. These findings are consistent with numerous reports describing filamentous thermophilic bacteria, most notably cultivated members of the sister phylum *Ignavibacteria* that range in length from 1 µm to greater than 15 µm^{24,25}.

Unique CRISPR-Cas fusion and limited biogeographic distribution

CRISPR (clustered regularly interspaced short palindromic repeats) elements and *cas* (CRISPR-associated) genes across the *Ca.* Kryptonia genomes were recovered, and are suggestive of defense against viral attack. A novel fusion between two different CRISPR-Cas types (type I and III; subtypes I-B and III-A) was identified in all genomes. This unusual fusion contained the full gene set for components responsible for the multistep CRISPR processes for spacer acquisition, CRISPR locus transcription and maturation, and final nucleic acid interference^{26,27} (Supplementary Fig. 5). This observation represents the first report of a type I-B/type III-A CRISPR-Cas

fusion and expands the known genetic diversity of CRISPR-Cas loci. Based on reconstruction of repeat-spacer arrays, the *Ca*. Kryptonium thompsoni genomes appear to represent a clonal CRISPR population without active spacer acquisition, while the *Ca*. Kryptobacter tengchongensis genomes are considerably dynamic in terms of a mosaic spacer collection (Supplementary Text; Supplementary Tables 6 and 7). These findings suggest that the CRISPR-Cas encoded by *Ca*. Kryptobacter tengchongensis is highly active, while the *Ca*. Kryptonium thompsoni genomes are not actively acquiring spacers through the CRISPR-Cas system.

To verify the limited biogeographic distribution of *Ca*. Kryptonia, we systematically surveyed the collection of 640 Gb of assembled metagenomic data from 4,290 environmental samples (including 169 samples from geothermal springs and hydrothermal vents) for the presence of a genomic signature beyond our initial search using SSU rRNA fragments from 100 kbp contigs (Fig. 3; Supplementary Table 8). Further, we searched against all available SSU rRNA data from the SILVA database¹⁶ for additional *Ca*. Kryptonia phylotypes and did not recover a highly similar match. Using this expanded search, we found evidence for *Ca*. Kryptonia in a total of twenty metagenomes, which included only three additional geographic sites compared to our initial SSU rRNA survey (Supplementary Table 8). The environments where this phylum was found were similar to the settings where we first discovered the genomic presence of *Ca*. Kryptonia: all were high-temperature (≥ 70°C), pH-neutral (6.4 – 8.0) settings. In sum, the limited range of *Ca*. Kryptonia is reflected in the observation that genomic signatures were found in nine unique geographical locations from a total of twenty-three

pH neutral hot springs currently sampled by metagenomics, and absent from the 1,614 unique locations represented by 4,290 metagenomic samples.

Additional metagenomic searches specific for all CRISPR repeat-spacer arrays collected from the Ca. Kryptonia genomes resulted in a similar pattern of limited biogeographic distribution (Fig. 3; Supplementary Table 7). We identified shared spacers across Ca. Kryptonia populations in geographically distinct geothermal springs. For example, shared spacers were identified between the Ca. Kryptobacter tengchongensis JGI-2 and JGI-3 genomes despite sampling from separate geothermal pools in China. Further, shared spacers were identified across exceptionally wide geographic distances including Canada and Nevada (Ca. Kryptonium thompsoni JGI-4 and the Great Boiling Springs metagenome), and China and Nevada (Ca. Kryptobacter tengchongensis JGI-2 and the Great Boiling Springs metagenome) (Fig. 3). Remarkably, we also found spacer matches to a set of metagenomic contigs that we assigned as viral due to their linkage to known viral genes, from these same samples and metagenome samples collected from Yellowstone National Park²⁸ (Fig. 3; Supplementary Text; Supplementary Fig. 5; Supplementary Table 9). These genomic recruitment and spacer signature data suggest that Ca. Kryptonia is present in additional geothermal spring sites and that viruses which appear to infect Ca. Kryptonia circulate across wide geographic space as revealed from the conserved infection vestiges.

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Metabolic potential of 'Candidatus Kryptonia'

The availability of multiple nearly complete *Ca.* Kryptonia genomes from both GFMs and SAGs enabled metabolic and putative functional predictions for this novel candidate phylum, as well as insights into some of the unique properties and notable absence of function for the individual genera. Approximately 50% of the predicted composite proteome for the Ca. Kryptonia genomes showed similarity to a diverse array of FCB superphylum members, with 11.3% and 1.96% best matches to thermophilic members of the phylum *Ignavibacteria* and *Caldithrix abyssi*, respectively (Supplementary Fig. 6). The conserved Por secretion system C-terminal sorting domain (TIGR04183), found exclusively in members of the FCB superphylum⁹, was recovered in all GFMs and SAGs, and altogether totaled 811 predicted proteins across the Ca. Kryptonia genomes. Reverse gyrase, the presumptive gene indicator for the extreme thermophilic and hyperthermophilic lifestyle in bacteria and archaea²⁹, was found in all Ca. Kryptonia genomes, which suggests that most, if not all members, of this lineage are extreme thermophiles or hyperthermophiles. Further, we found evidence for horizontal gene transfer of the reverse gyrase from the crenarchaeal order Thermoproteales (Supplementary Text; Supplementary Fig. 7) and hypothesize that Ca. Kryptonia's thermophilic traits might have been acquired via lateral gene transfer rather than ancestral inheritance.

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Ca. Kryptonia is a motile heterotroph with a complete tricarboxylic acid cycle and key metabolic enzymes for Embden-Meyerhof glycolysis and the pentose phosphate pathway. We found evidence for a complex oxidative phosphorylation pathway, which points towards aerobic respiration (Fig. 4; Supplementary Table 10). An elaborate and unique respiratory pathway for the redox transformation of iron is encoded in the Ca.

Kryptonia genomes with similar, yet non-homologous components to the well-characterized Mtr-like respiratory pathway³⁰ (Supplementary Fig. 8). Altogether, *Ca.* Kryptonia has the machinery to carry out ferric iron respiration under thermophilic conditions and likely vies with archaeal community members to impact metal biogeochemistry in these geothermal springs.

Ca. Kryptonia hosts the genomic potential for aromatic hydrocarbon degradation via oxidation to catechol, and subsequent catechol meta-cleavage (Fig. 4). Further, the Ca. Kryptonium thompsoni genomes encode a putative gene complement for the anaerobic degradation of aromatic amino acids or similar compounds, notably represented by a phenylacetyl-CoA oxidoreductase homologous to the hyperthermophilic archaeon Ferroglobus placidus³¹. This feature appears to be the first example of an extremely thermophilic or hyperthermophilic bacterium with the presumptive capacity to completely mineralize aromatic compounds, and holds biotechnological potential as well as implications for carbon cycling within geothermal springs³².

Unexpected metabolic deficiencies identified in Ca. Kryptonia

An unexpected observation was that all *Ca.* Kryptonia genomes had conspicuous nutritional deficiencies, displaying gene loss for many biosynthetic pathways, including thiamine, biotin and amino acids, such as the evolutionarily conserved histidine biosynthesis³³ (Fig. 4; Supplementary Table 10). While obligately host-dependent microbes and some free-living organisms with reduced genomes are known to omit a suite of anabolic pathways^{34,35}, the *Ca.* Kryptonia genomes do not appear to have

signatures of either lifestyle. An analysis of 759 high-quality FCB superphylum genomes indicate the near-complete *Ca.* Kryptonia genomes are distinct from free-living microbes in terms of amino acid pathway coverage and genome size, yet are not highly reduced compared to obligate symbionts (Supplementary Fig. 9). These findings suggest that *Ca.* Kryptonia has potentially evolved functional dependency on other microbes in order to acquire necessary metabolic requirements.

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To explore the existence of possible microbial partners, we performed a cooccurrence analysis of SSU rRNA sequences retrieved through their targeted assembly from an expanded set of 22 geothermal springs metagenomes (Supplementary Text; Supplementary Table 11). An analysis of co-occurrence patterns for clusters of taxonomically coherent groups (clustered at 90% sequence identity) revealed a subset of taxonomically clustered groups (phylotypes) highly correlated with the abundance of Ca. Kryptonia (Supplementary Table 12). These clusters included an Armatimonadetes lineage, which had the highest correlation value, three separate lineages of *Chloroflexi*, and *Thermus* spp. (Fig. 5). For the twelve metagenomes in which *Ca.* Kryptonia's SSU rRNA was reconstructed, the *Armatimonadetes* lineage was found to co-occur in seven of those metagenomes at similar sequence coverage to the Ca. Kryptonia genomes, and was conspicuously absent across all other metagenomes surveyed. To explore the potential of the Armatimonadetes lineage to complement the metabolic deficiencies identified in Ca. Kryptonia, we reconstructed three nearly complete genomes of Armatimonadetes (Fig. 2; Supplementary Tables 3 and 13) to infer metabolic potential and signatures of possible metabolic exchange and interaction. Analysis of the reconstructed genomes identified metabolic features complementary to those of Ca.

Kryptonia, such as histidine, cysteine and methionine, proline, aspartic acid, and thiamine biosynthesis, and degradation of pentoses (Fig. 5B; Supplementary Text; Supplementary Table 13). Furthermore, in the reconstructed *Armatimonadetes* genomes we also identified a CsqG family protein, which forms transmembrane channels for secretion of "functional amyloids," a class of bacterial proteins capable of assembling highly stable fibers through a nucleation-precipitation mechanism³⁶. "Functional amyloids" play major roles in adhesion to surfaces and biofilm formation in diverse bacteria including Escherichia coli, Caulobacter crescentus and Bacillus subtilis³⁷. Further, the CsqG-like transporter was located in a six-gene conserved cluster containing a predicted subtilase-family peptidase and a putative secreted protein with four copies of a "carboxypeptidase regulatory-like domain" (Pfam13620) (Supplementary Fig. 10). This domain is a member of the transthyretin clan and has been found to form amyloid in physiological conditions³⁸. We hypothesize that this cluster in the *Armatimonadetes* genomes encodes for synthesis, secretion and assembly of "functional amyloid," in which other members of the community may be embedded. On the other hand, the Ca. Kryptonia genomes encode many proteases and peptidases, which may be responsible for remodeling and digestion of this extracellular matrix.

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Other co-occurring lineages with *Ca.* Kryptonia include the *Thermus* spp. cluster (Supplementary Table 12). Interestingly, Ca. Kryptonia might complement an incomplete denitrification pathway in *Thermus* spp., which may be responsible for high rates of nitrous oxide production^{39,40}. *Thermus* spp. have been experimentally characterized to reduce nitrate to nitrous oxide but lack the capacity to subsequently

produce dinitrogen^{39,40}. Kryptonia encodes a nitrous oxide reductase (EC 1.7.2.4) but lacks other components of the denitrification pathway (Supplementary Text; Supplementary Table 14). Taken together, we hypothesize that *Ca.* Kryptonia may participate in a partnership with other organisms, such as the *Armatimonadetes*, or might interact with a broader consortium of microbes within the geothermal spring environment.

Discussion

A comprehensive survey of a global set of assembled metagenomic data for novel microbial lineages has resulted in the discovery of a new bacterial candidate phylum in geothermal springs. The high-quality draft genome assemblies enabled by complementary approaches from metagenomic data and single-cell genomics data for *Ca.* Kryptonia facilitated delineation of the host-virus interaction across geographically distant sites. Further, we observed a novel fusion between two different CRISPR-Cas types, representing the first report of a type I-B/type III-A CRISPR-Cas fusion and expanded the known genetic diversity of CRISPR-Cas loci.

The metabolic capacity for *Ca.* Kryptonia provides evidence for a unique heterotrophic lifestyle with the putative capacity for iron respiration within a consistent ecological niche in geothermal springs. An unexpected observation was that all *Ca.* Kryptonia genomes had conspicuous nutritional deficiencies, which led to the hypothesis of a microbial partnership or interaction with a broader consortium of microbes. Subsequent genome reconstruction of genomes from a co-occurring *Armatimonadetes* lineage indicated potential complementarity for those metabolic

features presumably absent in *Ca*. Kryptonia. It is well recognized that certain marine microbes, such as SAR11 (ref.⁴¹) and SAR86 (ref.⁴²), lack a variety of anabolic pathways and likely rely on other microbial community members to supplement their requirements. Within geothermal springs, the growth of chlorophototroph *Candidatus* Chloracidobacterium thermophilum in the laboratory was shown to depend upon two heterotrophs, *Anoxybacillus* and *Meiothermus* spp., due to lack of biosynthetic pathways for branched-chain amino acids, lysine and cobalamin⁴³. Our study suggests that dependency on other organisms within the geothermal spring community might be a more common occurrence than previously appreciated, perhaps contributing to challenges in obtaining many of these lineages as isolated monocultures. Future efforts to delineate this hypothesized interaction, particularly utilizing microscopy methods to visualize these uncultivated cells *in situ*, will further contribute to our understanding of *Ca*. Kryptonia and its role within the environment.

Geothermal springs have been heavily surveyed as a rich source of novel microbial branches on the tree of life^{18,44}, yet our results indicate that additional phylogenetic novelty has yet to be captured from these environments. The discovery of a new candidate phylum emphasize that extraordinary microbial novelty is likely still awaiting discovery using the vast metagenomic data assembled from locations sampled globally.

344 345 346 347 Materials and methods 348 Metagenomes 349 All publicly available metagenome datasets from IMG/M were used in the study¹⁴. The 350 metagenomes can be accessed at http://img.jgi.doe.gov and associated metadata can 351 be found in the GOLD database at http://genomesonline.org. 352 Metagenomic binning 353 Tetranucleotide-based binning methods were implemented as previously described to 354 recover near-complete genomes from metagenomes⁴⁵. Both single metagenomes and 355 combined metagenome assemblies were used to recruit additional contigs that 356 harbored the same tetranucleotide signature, and the raw reads were subsequently re-357 assembled using SPAdes version 3.1.0 (ref. 46). 358 SAG generation 359 Sediment samples were collected from Dewar Creek hot spring (49.9543667°, -360 116.5155000°) near the source of the hot spring on September 28, 2012, from the Jinze 361 pool (25.44138°, 98.46004°) on August 12, 2012, and from the Gongxiaoshe pool 362 (25.44012°, 98.44081°) on August 9, 2011. Samples were mixed with glycerol-TE buffer 363 for cryopreservation and stored at -80°C. Single cells were isolated using FACS, lysed, 364 and subjected to whole-genome amplification (WGA) as previously described with the 365 following modifications: the alkaline lysis was preceded by a 20 min digest with 366 lysozyme (Epicentre) at 30°C; WGA was performed with a REPLI-g Single Cell Kit

(Qiagen) with a scaled-down reaction volume of 2 µl; and the amplification reaction was incubated for 6 hr at 30°C. WGA reactions were diluted 10-fold, then aliquots were further diluted 200-fold for PCR screening targeting the V6-V8 regions of the SSU rRNA using a QuantiNova SYBR Green PCR kit (Qiagen) for 45 cycles of amplification⁹. PCR products were purified and sequenced, and SAGs matching Kryptonia SSU rRNA sequences were selected for shotgun sequencing.

SAG sequencing, assembly and QC

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Draft genomes for the eighteen SAGs were generated at the DOE Joint Genome Institute (JGI) using the Illumina MiSeq technology according to standard protocols (http://www.jgi.doe.gov/). Assembly was performed using SPAdes version 3.1.0 (ref. 46) using the --sc flag to denote MDA-derived data to account for uneven coverage of the single-cell genomes. Quality control and contaminant removal from the resultant assemblies was achieved using a two-step process. First, all assembled reads were used as input for a newly developed single-cell decontamination method (ProDeGe)⁴⁷, which uses both taxonomic and k-mer based decisions to flag putative non-target contigs. Since the taxonomic information was limited to phylum-level designations, we further supplemented this procedure with direct mapping to the genomes reconstructed from metagenomic data. For mapping, a combination of blast and blat were implemented to validate correct recruitment of the assembled SAG contigs to Ca. Kryptonia-specific GFM scaffolds. This method was important for retaining CRISPR/Cas genetic regions since ProDeGe had the tendency to flag these contigs based on divergent k-mer frequencies. Gene annotation was performed within the Integrated Microbial Genomes (IMG) platform developed by the DOE Joint Genome Institute¹⁴.

SSU rRNA phylogeny

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Full-length SSU rRNA gene sequences from Ca. Kryptonia were aligned using the SINA aligner¹⁵ to a comprehensive database of references (SILVA-NR version 119)¹⁶. A total of 187 full-length bacterial and archaeal reference sequences were selected from the SILVA database, and 1,354 distinct alignment patterns were used, and filtered using the E. coli positional mask. A maximum likelihood tree was calculated from the masked alignments with 100 bootstrap resamplings using the Generalized Time-Reversible model with G+I options in RAxML version 7.6.3 (raxmIHPC-PTHREADS-SSE3 -f a -x 12345 -p 12345 -# 100 -T 5 -m GTRGAMMAI)48. To resolve placement within the FCB superphylum, a subset of 77 FCB superphylum members and 37 archaeal references sequences were selected and phylogenies constructed using two separate algorithms: the GTR+G+I model (used with maximum likelihood, RAxML⁴⁸) and Bayesian inference (MrBayes⁴⁹). Node stability was evaluated using a rapid bootstrapping analysis (RAxML, 100 runs) and posterior probabilities (MrBayes, 2.4 million generations, burnin of 25%). Alignments and phylogenetic trees are available at http://portal.nersc.gov/dna/microbial/prokpubs/Kryptonia. Catalyzed Reporter Deposition Fluorescence In Situ Hybridization (CARD-FISH) An oligonucleotide probe specific for Ca. Kryptonia (Kryp56; 5'-CCGTGTCCCTGACTTGCA-3') was designed in ARB (version 6.0.2)⁵⁰. The probe is a perfect match to 19 of the 22 Ca. Kryptonia SSU rRNA gene sequences recovered in this study, and contains two or more mismatches to all SSU rRNA gene sequences in the SILVA-NR database (version 123)¹⁶. The probe sequence was synthesized by Biomers.net (Ulm, Germany) with horseradish peroxidase (HRP) conjugated to the 5'

end. Cells from Dewar Creek sediment were separated from particulates by brief vortexing followed by centrifugation. Suspended cells were preserved with dimethyl sulfoxide (DMSO) and stored at -80°C. The cells were permeabilized with lysozyme and CARD-FISH was performed based on the protocol of Pernthaler and colleagues⁵¹. Hybridization was carried out at 46°C with 20% formamide, and the amplification was performed with tyramides conjugated to Alexa 488 (Life Technologies, #T20948). The optimal formamide concentration and specificity was predicted using mathFISH⁵² and the DECIPHER ProbeMelt tool⁵³ (Supplementary Table 15), and confirmed empirically by performing CARD-FISH on the Dewar Creek cells over a gradient of formamide concentrations (10 – 35%). Samples were counterstained with 4',6-diamidino-2phenylindole (DAPI) in VECTASHEILD Antifade Mounting Media (Vector Laboratories, #H-1200). Cells were visualized and imaged using a Leica DM6000B microscope using a HCX PL APO 100X oil immersion objective. Conserved single-copy and housekeeping gene phylogenetic inference A set of 56 universally conserved single copy proteins in the Bacteria and Archaea was used for phylogenetic inference (Supplementary Table 16). Marker genes were detected and aligned with hmmsearch and hmmalign included in HMMER3 (ref. 54) using HMM profiles obtained from phylosift (http://phylosift.wordpress.com/)⁵⁵. Alignments were concatenated and filtered⁵⁶. Housekeeping genes were aligned using MAFFT with mafft-linsi option⁵⁷. Best substitution model was selected using prottest⁵⁸. Phylogeny was inferred using maximum likelihood methods with RAxML (version 7.6.3)⁴⁸. Tree topologies were tested for robustness using 100 bootstrap replicates with the LG+I+G model (raxmlHPC-PTHREADS-SSE3 -f a -x 12345 -p 12345 -# 100 -m

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436 PROTGAMMALG -T 5). Trees were visualized using Dendroscope⁵⁹. The concatenated 437 protein alignment and phylogenetic tree is available at 438 http://portal.nersc.gov/dna/microbial/prokpubs/Kryptonia. 439 Phylogenetic distribution of predicted proteins 440 The taxonomic distribution of all proteins across the genomes reconstructed from 441 metagenomic data along with the Ca. Kryptonia SAGs was compiled based on best 442 matches to a comprehensive protein database of high-quality non-redundant bacterial 443 and archaeal isolate genomes¹⁴. This search was performed using usearch (version 444 $(7.0)^{60}$, where a protein match was considered for proteins with $\geq 30\%$ sequence identity 445 across ≥ 50% of the query alignment length. Phylogenetic affiliation at the phylum level 446 was assigned for top matches, while proteins lacking a match according to the above 447 criteria were noted as 'no match.' 448 Biogeographic survey of Ca. Kryptonia against available metagenomic data 449 All genomic data for Ca. Kryptonia was searched against the assembled metagenomic 450 data from 4,290 environmental samples using blat with the -fastMap option⁶¹. Significant 451 matches for non-ribosomal genomic regions were considered for sequences ≥ 250 bp in 452 length and with ≥ 75% identity threshold. For metagenomic contigs mapping to the 453 ribosomal operon, a 97% identity threshold was used to capture only high-quality 454 matches to Ca. Kryptonia. Visualization of metagenomic matches globally was 455 performed using the R package 'maps'62. All genomic matches can be found in

CRISPR-Cas locus type determination

Supplementary Table 8.

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We used 99 CRISPR-associated (*cas*) gene sequence alignments and hidden Markov models from the TIGRFAM database (originally built by Haft *et al*.⁶³ and later expanded by Zhang *et al*.⁶⁴) to precisely find and identify Cas family members within the scaffolds of the *Ca*. Kryptonia genomes. We recovered and classified the corresponding CRISPR type for complete and partial CRISPR-Cas loci in all genomes following the unified CRISPR classification from 2011 (ref.⁶⁵).

CRISPR repeat-spacer arrays analysis

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The CRISPR Recognition Tool (CRT)⁶⁶ was used to detect CRISPR repeat-spacer regions across all Ca. Kryptonia assembled scaffolds using specific parameters (according to the JGI's annotation pipeline). In the case of Ca. Thermokryptus mobilis GFM JGI-1, we were unable to detect spacers, and therefore we additionally used the CRISPR assembler algorithm (Crass)⁶⁷ on the raw reads. Spacers were manually curated to cull false positives from the dataset that clearly did not represent authentic spacer regions (in sum, 38 false positives). Potentially active repeat-spacer arrays were inferred based on direct association with a cas gene locus. We also considered the isolated repeat-spacers arrays when they shared the same repeat sequence with associated cas genes. CRISPRmap^{68,69} was used to further characterize identified repeat regions. From a total of 1,031 trusted spacers, we next clustered these into 795 groups based on identity ≥ 90% over the whole spacer length. Spacer groups were BLAST queried against distinct databases including Ca. Kryptonia genomes, reference public plasmid and viral datasets (from NCBI), and across the broad available metagenomic space (IMG/M).

Extraction, assembly and clustering of SSU rRNA genes from shotgun

metagenomic data

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Raw reads aligning to 16S and 18S rRNAs were collected for 22 metagenomes (Supplementary Table 11) from geothermal environments using hmmalign⁵⁴ against hmm models representing bacterial, archaeal and eukaryotic sequences⁷⁰ and also by BBmap with default settings⁷¹ against sequences from the SILVA database (version 119)¹⁶ dereplicated at 95% identity using UCLUST⁶⁰. Collected paired-end Illumina reads were merged using BBmerge⁷¹ and assembled using Newbler (v. 2.8)⁷² with -ml 60 -mi 99 -rip options. Resulting contigs and scaffolds were screened using cmalign from Infernal 1.1 package⁷³ and Rfam 16S and 18S rRNA models (RF00177.cm, RF01959.cm and RF01960.cm)⁷⁴. 16S and 18S rRNA sequences longer than 300 nt were retained and trimmed using cmalign against the best-matching model with '-matchonly' option to remove introns. Reference sequences from the SILVA database were trimmed using cmalign with a domain-specific model and '--matchonly' option, and clustered together with 16S sequences extracted from shotgun metagenome data using UCLUST and percent identity cutoffs of 94%, 92% and 90%. Clusters including sequences from at least two metagenome samples were retained and their abundances in metagenome samples were computed by multiplying the length of SSU rRNA sequence by the average coverage. Taxonomy was assigned to the clusters as last common ancestor (LCA) of SILVA reference sequences included in the cluster, or as LCA of SILVA sequences in the larger cluster obtained by co-clustering SILVA and metagenome sequences at 83% identity. Spearman's rank-order correlation of cluster abundances was used to estimate co-occurrence of the clusters in metagenome data.

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Accession Codes

- 505 Genome sequence data have been deposited as Whole Genome Shotgun projects at
- 506 DDBJ/EMBL/GenBank with the accession codes LDNR00000000, LDYP00000000-
- 507 LDZE00000000, and LDXU00000000-LDYB00000000. The versions described in this
- paper are versions 01000000. The annotated genomes and single-cell assemblies can
- be accessed via IMG (http://img.jgi.doe.gov).

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Author Contributions

- 719 E.A.E-F., N.C.K., and N.N.I. designed the project; P.F.D., B.P.H., S.E.G., A.L.B., H.D.,
- 720 B.R.B., and W-J.L. provided the samples; J.J., D.G., R.M., and T.W. performed the
- single-cell experiments; A.E.D. and J.P-R. performed the CARD-FISH experiments;
- 722 E.A.E-F., D.P-E., J.J., P.F.D., A.P., T.W., N.C.K., and N.N.I. analyzed the data; E.A.E-
- 723 F., D.P-E., N.C.K, and N.N.I. wrote the manuscript with significant input from P.F.D.,
- 724 B.P.H., T.W., and E.M.R. All authors discussed the results and commented on the
- 725 manuscript.

Competing interests The authors declare no competing interests. Figure 1. New lineage identified using metagenomic and single-cell genomic approaches. Workflow used to (A) identify novel SSU rRNA gene sequences globally, along with (B) single-cell genomics pipeline to screen and sequence single cells isolated from geothermal springs samples. For the three geothermal spring environments, we sequenced 13, 2, and 3 SAGs, respectively. Figure 2. Maximum likelihood concatenated protein phylogeny and cell imaging for Ca. Kryptonia. (A) Phylogeny was based on concatenation of 56 conserved marker proteins, where at least 10 marker proteins were used to infer SAG phylogenetic placement (with the exception of JGI-22 with only six marker proteins recovered). Bootstrap support values ≥ 50% are shown with small circles on nodes with robust

phylogenetic support. The *Fibrobacteres-Chlorobi-Bacteroidetes* (FCB) superphylum is shown in the gray shaded region. Expanded phylogenetic tree for *Ca.* Kryptonia shows the placement of the proposed four genera represented by GFMs and SAGs, along with the estimated genome completeness shown in parentheses. (**B**) A *Ca.* Kryptonia-specific FISH (fluorescence *in situ* hybridization) probe was designed and used to visualize cells from Dewar Creek Spring sediment samples. *Ca.* Kryptonia cells hybridizing with the probe are green, while other cells are visualized with 4',6-diamidino-2-phenylindole (DAPI; blue). Scale bar, 5 μm.

Figure 3. Limited, yet widely dispersed biogeographic distribution of *Ca*.

Kryptonia genomes and CRISPR spacers. All genomic content from the *Ca*.

Kryptonia GFMs and SAGs was used to comprehensively search the collection of 640
Gb of assembled metagenomic data from 4,290 environmental samples, including 169
samples from geothermal springs and hydrothermal vents denoted by red triangles
(temperature ≥ 50°C). Marked circles are as follows: (A) Great Boiling Spring,
Nevada^{20,21}, (B) Dewar Creek Spring, Canada¹⁹, (C) Jinze pool, Yunnan Province,
China²², and (D) Gongxiaoshe pool, Yunnan Province, China²². Significant matches
were determined for sequences ≥ 250 bp in length and with ≥ 75% identity threshold for
non-ribosomal genomic regions. For metagenomic contigs mapping to the *Ca*. Kryptonia
ribosomal operon, a 97% identity threshold was used to capture only high-quality
matches to *Ca*. Kryptonia. For CRISPR spacers, only significant matches allowing for
up to 3 bp mismatch along the entire length of the spacer were considered. The *Ca*.

Kryptonia genomic hits can be found in Supplementary Table 8 and the manually curated spacer hits can be found in Supplementary Table 7. Figure 4. Reconstructed metabolic capacity of Ca. Kryptonia. Key metabolic predictions and novel features identified in Ca. Kryptonia GFM and SAGs, with full gene information available in Supplementary Table 10. **Figure 5.** Co-occurrence patterns and metabolic complementarity with *Ca.* Kryptonia. (A) Spearman-rank correlation values were calculated based on reconstructed SSU rRNA sequences across 22 geothermal spring metagenomes, and led to the identification of a cluster of highly correlated phylotypes with Ca. Kryptonia. Armatimonadetes (cluster 3107) had the highest correlation value ($\rho = 0.82$) with Ca. Kryptonia. (B) Biosynthetic pathways present in the *Armatimonadetes* genome which complement missing components in Ca. Kryptonia. Each arrow represents an enzymatic component of the biosynthetic pathways; arrows highlighted in blue are contributed by the Armatimonadetes, while arrows highlighted in dark orange are

contributed by Ca. Kryptonia. Black arrows indicate enzyme was not recovered in either.